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- Tackified pressure-sensitive adhesive.
- This invertion provides tackfiled acrylic ultraviolet-radiation polymerized pressure-sensitive adhesives having improved achesive properties. The adhesives contain an acrylic ultraviolet-radiation photopolymerized polymer having an alkyl acrylate monomer, the alkyl groups of which have an average of 4 to 12 carbon atoms, a monoethylenically unsaturated strongly polar copolymerizable monomer, and a hydrogenated rosin ester tackfying agent. The achesive has comparable adhesion to solvent-based tackfied acrylic pressure-sensitive adhesives containing to 5 of times as much tackfying agent.

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# TACKIFIED PRESSURE-SENSITIVE ADHESIVE

## Background of the Invention

#### Field of the Invention

This invention relates to photopolymerized ecrytic pressure-sensitive adhesives, and adhesive tackified with certain tackifying resins.

## Description of the Related Art

Acrylate pressure-sensitive adheaves are well known in the art. In U.S. Patent No. Re 24.906 (Uhrich), sityl acrylate copolymers are described which are predominately alkyl esters of acrylic acid having from 4 to 14 carbon stioms, and further comprise a minor amount of a poter copolymerizable monomer such as acrylic acid. Such adheaves are widely oppular es they are readily available and provide a good balance of tack, shear and peel properties at a relatively flow cost.

U.S. Patent No. 4,161,752 (Martens et al) discloses a process for ultraviolet photopolymerization of alkyl acrylate seters and polar copolymerizable monomers to form the acrylate copolymer. Martens teaches that intensity and spectral distribution of the irradiation must be controlled in order to attain destrable cohestive strengths and peel resistance. The photopolymerization is preferably carried out in an inext atmosphere as oxygen tends to inhibit the reaction.

Additional patents disclose ultraviolet radiation of acrylate adhesives. U.S. Patent No. 4,384,972 (Moon) discloses the use of N-vinylpyrrolidone as the polar copolymerizable monomer in the acrylate adhesive copolymer to provide a pressure-sensitive adhesive with high adhesion to automotive paints. U.S. Patent No. 4,391,807 (Vesley) discloses the use of specific chromophore- substituted-halomethys-trizines as photoactive crossifixers for carylate copolymers. U.S. Patent No. 4,599,265 (Esmay) discloses a readify pressure-sensitive achieves the provided provided to the polar polar provided provided

The above-cited Moon patent, which concerns pressure-sensitive adhesives designed specifically to provide enhanced adhesion to automotive paints, teaches that teachings can be blended into the photoactive mixtures of monomers from which those pressure-ansitive achesives are photopolymerized, but warns, at 5 column 6, lines 3-12, that "the addition of any such material adds complexity and hence expense to an otherwise simple, straight forward, economical process, and is not preferred, except to achieve specific photopolymerizable mixture of monomers other interferes with the photopolymerization and prevents the attainment of the desired adhesive and cohesive properties.

U.S. Patent No. 4,243,500 (Glennon) discloses a pressure-sensitive adhesive formed from a composition comprising at least one monofunctional unsaturated acrylate ester monomer, essentially saturated tackifying rest polymers, non-crystallizing elastometic material, and an initiator responsive to ultraviolet light, or other penetrating radiation. Glennon specifies the use of UV light within a wavelength range of the lamps disclosed in Martens et al.

Glennon states that the tackifying reain may be a substance or mixture of substances selected from the group consisting of seters of rosis, hydrogenated esters of rosis, modified rosin esters, esters of polymerized rosin, esters of hydrogenatic rosis, hydrocarbon rosis, flush monopolymers of alpha-methyl styrene, sophere terpene hydrocarbon rosis, aromatic modified CS hydrocarbon resis, dispha-methyl styrene copolymer resis, beit-pinene terpene resins, polycyclic hydrocarbon rosis, and technical hydrocalbely alcohol. However, many of these essentially saturated resin polymers are unsuitable for use in the curing method of the above-roticed Martens patent due to incombility, which results in phase separation of the tackifying resin from the monomer mixture, excessive UV absorption which retards the photopolymerization of the monomers is impeded. The Glennon adhesive requires the use of from about 50 parts to about 250 parts to

tackifying resin per hundred parts acrylate ester monomer in order to obtain a satisfactory adhesion level.

Such levels of tackifler increase the glass transition temperature and act as chain transfer apents during the polymerization. This results in the pressure-sensitive adhesives having lowered molecular weights and reduced orbiseive strengths. Further, many tackifiers tend to migrate to the surface of the adhesive upon

a aging, deteriorating the adhesion.

U.S. Patent No. 4,50,883 (Hot et al) discloses a pressure-sensitive adhesive composition containing as a polymer component, an addition-polymerization polymer of an acryl based polymer having sticking properties at room temperature and one or more ethylenically unsaturated monomers capable of forming a homopolymer or copolymer having a glass transition point of at least 273 K. The addition-polymerization polymer is prepared by polymerizating one or more ethylenically unsaturated monomers in the presence of the acryl-based polymer by solution polymerization or bulk polymerization using redical polymerization catalysts, but polymerization is initiated by energy in the form of light, etc. Miscollaneous additives such as coloring agents, fillers, arti-éging agents, and tacktifiers are disclosed as possible additions.

U.S. Petern No. 4.418.120 (Kealy et al.) discloses a pressure-sensitive adhesive tape which is made by U.S. Petern No. 4.418.120 (Kealy et al.) discloses a pressure-sensitive adhesive tape which is made by coating a sheet backing with a solution of isocityl scrylate: acrylic acid copplymer containing a tackifying roslin ester and an antiboxidant, evaporating the solvent, and crosslinking the adhesive U.S. Patent 48.45.711, (Winslow et al.) discloses a removable pressure-sensitive adhesive tape, the adhesive law which is an emulsion polymerized copplymer of allyd scrylate such as isocotyle acrylate and a small amount of emulsifier monomer and a tackfying reains selected from hydrogenated rosin esters, polytorpene, of emulsifier monomer and a tackfying reains selected from hydrogenated rosin esters, polytorpene, or polymerized allyl styrene, and polymerized petroleum derived monomer resists. Most of these resins are unsuitable for use in in thus photopolymerized acrylate pressure-sensitive adhesives.

US. Patent 3,888,247 (Slenvall) discloses a first aid bandage having an achesive layer applied over a flexible, preferably microporous backing. The adhesive layer is an acrylate type pressure-sensitive adhesive adsorbed in Utrich, in a 946 ratio, and contains 40 person of a highly stabilized ester resin tackfler, as described in Utrich, in a 946 ratio, and contains 40 person of a highly stabilized ester resin tackfler, as commercially available from Hercules Chemical Co. under the trade name Foral 85, U.S. Patent 4,335,171 (Zent) discloses a pressure-sensitive adhesive transfer tape in which the achieve is a laminate of two gravity disciplinations. The first lamina is stated to be a soft pressure-sensitive adhesive consisting physically disaminal naminaes. The first lamina is stated to be a soft pressure-sensitive adhesive consisting essentially of an isocotyl acrylateacrylic acid copolymer wherein the acid present is in the range of 3-12% essentially of an isocotyl acrylateacrylic acid copolymer wherein the acid present is in the range of 3-12% of the copolymer. The copolymer is tackifled with 50 parts of a hydrogenated resin ester or hydrogenated resin state or hydrogenated resin state or hydrogenated resin state or hydrogenated resin state or hydrogenated resin state.

The current inventor has discovered that use of a certain type of tackifying resin in small amounts results in adhesion equivalent to that of solvent polymerized adhesives containing 6 to 7 times as much tackifying resin.

# Summary of the invention

This invention provides tackified acrylic ultraviolet-radiation photopolymerized pressure-sensitive adhe-40 sives having improved adhesion to plastic surfaces. The adhesives of the invention have comparable adhesion to solvent-based tackified acrylic pressure-sensitive adhesives containing 6 to 7 times as much

tackifying agent. The adhesives comprise
a) from about 85 parts to about 98 parts of an acrylic ultraviolet-radiation polymerized polymer of

monomers containing:
(i) from about 93 to about 99 parts of an alkyl acrylate monomer, the alkyl groups of which have an

average of 4 to 12 carbon atoms, and (ii) correspondingly, from about 7 parts to about 1 part of a monoethylenically unsaturated strongly polar copolymerizable monomer,

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b) correspondingly, from about 15 parts to about 2 parts of a hydrogenated rosin ester tackifying agent, and c) from about 0.01 part to about 1 part of a photoinitator, based on the weight of the acrylic

c) from about UU1 part to about 1 part to a production of the facility of a production of the facility of the

As used in this application, the terms "tackfiler" "tackfilying resin", and "tackfilying agent" are all synonymous, and refer only to the hydrogenated rosin esters useful in the present invention, unless otherwise specifically stated.

All percentages, parts, and ratios herein are by weight unless otherwise specifically stated.

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### Detailed Description of the Invention

The tacklifled acrylic ultraviolet-radiation photopolymerized pressure-sensitive adhesive compositions of the present invention contain from about 2 parts to about 15 parts, preferably from about 5 parts bout 12 parts, or a tacklying agent selected from the group consisting of hydrogenated rosin esters. Rosin esters have a higher softening point than unmodified rosins, and higher molecular weight. Ethylene glyboo, glycerol, and pentaneythiol are the most connected to esterification. Rosin esters are quite stable, and resistant to hydrolysis, and such stability increases with hydrogenation. Surprisingly, earlylic ultraviolet-radiation photopolymerated pressure-sensitive adhesives tacklified with small amounts of hydrogenated rosin ester tacklifying agents have comparable adhesion to tacklifier. Such tacklified adhesives have better adhesion to plastics than do ultraviolet-polymerized acrylic ressure-sensitive adhesives to containing 8 to 7 times as much rosin ester adhesions containing no such tacklifier.

Preferred tackfying agents are highly hydrogenated. Such tackfilers, e.g., hydrogenated glycerine seters are commercially available from companies such as Hercules Inc., under the trade names Foral Pentalyn and Staybelfte, particularly, ForalTM 65, ForalTM 85, ForalTM AX, ForalTM 105, and StaybelfterTM Ester 10.

The acrylic utraviolar-radiation photopolymerized polymers (acrylic copolymer(s)) constitute from about 95 to about 98 parts of the pressure-sensitive adhesive compositions of the invention. Acrylic copolymers useful in achievive of the present invention are copolymers containing at least one alkyl acrylate monomer, have from 4 to about 14 carbon atoms. Included within this class of monomers are, for example, isooctyl acrylate, isonomyl acrylate, 2-ethyl-hexyl acrylate, decyl acrylate, bright a

Polarity, or hydrogen-bonding ability, is trequently described by the use of terms such as "poorly", or "moderately", and "strongly". References explaining the use of these and other subbility terms include "Solvents". Paint Testing Manual, 3rd Ed., Seward, G.G., Editor, American Society for Testing and Materials, Phila.Pa., 1972, and "A Three-Dimensional Approach to Solubility", Journal of Paint Technology, 38, No. 486, pp. 269-260.

The strongly polar copolymerizable monomers useful in the acrylic copolymers of the Invention may be acrylle acid, itaconic acid, hydroxyalkyl acrylates, cyanoslkyl acrylates, acrylamides or substituted acrylamides. While photopolymerizad acrylic copolymers using moderately polar monomers such as Nvinyl polar polar monomers acid as Nvinyl do not show the improved tack when tackfilled with low levels of hydrogenated rosin ester tackfilling agent as to the strongly polar monomers. The strongly polar copolymerizable monomer comprises up to about 7 parts, more preferably up to about 5 parts of the photopolymerizable polymer. Conventional pressure-sensitive adhesives containing strongly polar monomers in the low levels present in adhesives of the means the monomer comprises the property of the photopolymerizable polymer. Conventional pressure-sensitive adhesives containing strongly polar monomers in the low levels present in adhesives of the present invention.

The mixture of the photopolymerizable monomers and the tackifier also contains a photoinitiator to induce polymerizable of the monomers. Photoinitiators which are useful include the benzoin eithers such as benzoin methyl either or benzoin isopropyl either, substituted benzoin eithers such as anisole methyl either, substituted acetophenones such as 2.2-diethyoxyacetophenone and 2.2-dienthoxya-2-phenyl acetophenone, substituted alpha-ketols such as 2-diethyoxyacetophenone, aromatic sulfonyl chlorides such as 2-naphthalene sulfonyl chloride, and photoactive oximes such as 1-phenyl-1.1-proprediene-2-(0-ethoxy-carbonyl)-oxime. The photoinitiator is present in an amount of about 0.01 part to about 1 part by weight.

Where superior cohesive strengths are desired, the pressure-sensitive adhesive matrix of the novel multiacrystates such as 1,6-bxaraefold dearylate as well as those disclosed in U.S. Patent No. 4,379,201 (Heilmann et al.). Crosslinking is especially easy to control when photopolymeraizing the monomer in admittacrystate crosslinking agents to Cher types of crosslinking agents are also useful, e.g., any of the triazine crosslinking agents upon tin U.S. Patents 4,330,590 (Veslay), and 4,323,334 (Vesley et al.). Each of the crossfriking agents is useful in U.S. Patents 4,330,590 (Veslay), and 4,323,334 (Vesley et al.).

The instant invention also encompasses tapes containing at least one tackified acrylic ultraviolet-

radiation photopolymerized pressure-sensitive adhesive layer. Tapes of the invention may comprise more than one pressure-sensitive adhesive layer. In such tapes, the pressure-sensitive adhesive layers may comprise similar or different adhesives, in like or unlike thicknesses, having similar or different additives.

Where a foam-like pressure-sensitive adhesive tape is desirable, i.e., in those applications requiring a adhesion to low energy substrates such as polyethylene and polypropolyene, and coratin high solids automotive paint systems, a monomer bland comprising microbubbles may be used as a backing or core layer. The microbubbles may be glass, or they may be polymeric. The microbubbles should have an average diameter of 10 to 200 micrometers, and comprise from about 5 to about 55 volume percent of the pressure-sensitive adhesion layer. The thickness of the foam-like layer is preferable at leasts six times that 10 of each microbubble-free layer. When glass microbubbles are used, the thickness of the layer should exceed three times the average diameter of the microbubbles and twice the diameter of substantially every microbubble. When polymeric microbubbles are used, the thickness of the layer should be such that all of the microbubbles are substantially completely surrounded by the polymer matrix. The thickness of foam-like layers in preferred tapes of the livention range from 0.3 mm to about 4.0 mm in thickness.

years in presence uses or an emission and manufactured to the provided on a substantially. When a microbubble-free pressure-sensitive achiesive tape is desired to be provided on a substantially when a microbubble-free pressure-sensitive achiesive type in the tackfilled acrylic ultraviolet-radiation photopolymerized pressure-sensitive achiesive layer, (e.g., an alkylic acid acrylicte monomer and a strongly polar copolymerizable monomer with different ratios of the acrylic acid scriptate monomer and activatively, the polar sets of non-tertiary alcohol and at least one polar copolymerizable monomer. Alternatively, the polar monomer many also be selected from moderately polar monomers, such as N-vinyl pyrrolictone. N-vinyl monomer may also be selected from moderately polar monomer ms. Such layer may also comprise a capolicatam, acrylicitrile, vinyl chioride or dialityl phthalate. The preferred range of the polar monomer in such a layer ranges from 20% to about 50% of the total monomer mix. Such layer may also comprise a crosstrinking agent and other photopolymerizable ingredients including, but not limited to alkyl vinyl ethers, vinylidene chloride, styrene, and vinyl toluene, only in amounts that do not detract from the desired properties.

Other materials which can be blended with the polymerizable monomer mixture include plasticizers, reinforcing agents, dyes, pigments, fibers and fire retardants.

An especially useful filler material is hydrophobic silica as disclosed in U.S. Patent No. 4,710,536, (Klingen, et al.). In one preferred embodiment of the present invention, the pressure-sensitive adhesive layer (thingen, et al.) in one preferred embodiment of the present invention, the pressure-sensitive adhesive layer (thingen et al.) in one preferred embodiment of the present invention as a surface area of at least 10

The photopolymerizable compositions used in tapes of the invention are preferably prepared by premixing together the photopolymerizable monomers and photoinitiator, and for the tackfilled pressure-sensitive compositions, the tackfiler. This premix is then partially polymerized to a viscosity in the range of form about 500 cps to about 5,000 cps to achieve a coatable syrup. Alternatively, the monomers can be mixed with a thixotropic agent such as furned silica to achieve a coatable syrup composition.

This composition is coated onto a flexible carrier web and polymerized in an inert, i.e., oxygen-free, atmosphere, e.g., a nitrogen atmosphere. A sufficiently inert atmosphere can be achieved by covering a layer of the photoactive coating with a plastic film that is substantially transparent to ultraviolat radiation, and any irradiating through that film in air using lucrescent-type ultraviolat lamps which generally have an intensity of about 0.4 wattscm. If, instead of covering the polymerizable coating, the photopolymerization is to be carried out in an inert atmosphere, the permissible oxygen content of the inert atmosphere can be increased by mixing into the polymerizable monomer an oxidizable tin compound as taught in U.S. Patent Increased by mixing into the polymerizable monomer an oxidizable tin compound satisfy the U.S. Patent Increased by mixing into the polymerizable monomer an oxidizable tin compound satisfy the U.S. Patent Increased by mixing into the polymerizable monomer an oxidizable tin compound as taught in U.S. Patent Increased by mixing into the polymerizable monomer an oxidizable tin compound as taught in U.S. Patent Increased by mixing into the polymerizable monomer an oxidizable tin compound as taught in U.S. Patent Increased by mixing into the polymerizable monomer an oxidizable tin compound as taught in U.S. Patent Increased by mixing into the polymerizable monomer and oxidizable tin compound as taught in U.S. Patent Increased by mixing into the polymerizable monomer and increased by mixing into the po

The pressure-sensitive adhesive compositions may be coated onto a backing or substrate prior to polymerization. This may be accomplished using any conventional means such as roller coating, dip coating or extrusion coating.

As used herein, the term "tape" includes but is not limited to, those adhesive strips which are singlecoated adhesive layers permanently attached to a backing or support, double-coated adhesive strips having
flexible supports with an adhesive layer on both sides thereof, and adhesive strips with no support or
backing, such being typically releasably attached to a low-adhesion liner, and commonly called "transfer
tapes".

The following tests may be used to evaluate tapes of the invention.

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Static Shear

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Two flat, rigid stainless steel plates are bonded together by a piece of an adhesive layer 1.27 cm × 2.5 cm. Before testing, a 1000-g weight rests over the bonded area for 15 minutes. Then the bonded plates are placed in an air-circulating own which has been preheated to 70°C, and after 15 minutes, a 500-g weight is hung from one of the plates while the other plate is held in a position tilted 2° from the vertical to insure 5 against any peel forces. The time at which the weight falls is the "Static Shear Value at 70°C". If no failure occurs, the test is discontinued at 10,000 minutes. Only cohesive failures are reported.

#### Peel Adhesion Test

Peel adhesion tests were based on the method described in ASTM D903-49, "Peel or Stripping Strength of Adhesive Bonds."

#### 180° Peel

The adhesive layer to be tested is transferred onto 50 µm thick, chemically primed, aluminum foil, which then is sit to a width of 1.27 cm (1/2 inch). The resulting tape is self-adhered to a smooth stainless steel plate under the weight of a 2.04 kg hard-rubber-covered steel roller, 2 passes in each direction. After dwelling at 23° C for one hour, "180° Peel" is measured by moving the free end of the tape away from the Newtons por declimator (Nidtim), unloss otherwise noted.

### 90° Peel

The achiestre layer to be tested is transferred onto a 0.05-mm thick soft aluminum foll which then is sit to a width of 1.27 cm (1/2 inch). The resulting tape is self-adhered to a smooth stainless steel plate under the weight of a 5.5-kg hard-cubber-covered steel roller, one pass in each direction. After dwelling at 23 °C or 20 minutes, the "30" Peel" is measured by moving the free end of the tape away from the steel plate at 90 and at a rate of about 0.5 or per second using a tensite lester.

## Examples

## Example 1 and Comparative Example C1

A premix was prepared using 93.5 parts isooctyl acrylate, 6.5 parts acrylic acid and 0.04 parts irgacure 
651. This was partially polymerized by exposure to ultraviolet radiation to provide a coatable syrup having a 
10 parts of Foral 85 tackifying resin, 0.20 part of XL-353 (a triazine crossinter), and an additional 0.20 part 
Irgacure 651 was coated onto a polyethylene-coated paper to a uniform thickness of about 0.127 mm. 
Comparative Ezample 1 was prepared without tackifying resin. The surfaces of the polyethylene coated 
paper contacting the adhesive composition had low-adhesion release coatings on them.

These composites were exposed to ultraviolet radiation to promote polymerization of the syrup. The comparative peel adhesion is set forth in Table 1.

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Table 1

90° Peel Adhesion (N/dm.)		
Example ABS Polyethyler Plastic		Polyethylene
1 C1	70 55	55 22

# Examples 2-3 and Comparative Examples C2-C3

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A blend of 100 parts of the partially polymerized premix described in Example 1, 10 parts of tackifying resin Foral 85, 0.15 part of XL-330 (triazine crosslinker), and an additional 0.35 part of Irgacure 651 was coated onto a release coated polyethylene coated paper to a uniform thickness. Example 2 was coated to a thickness of about 0.051 mm. Example 3 was coated to a thickness of about 0.127 mm. These composites were exposed to ultraviolet radiation to promote polymerization of the syrup.

Comparative Examples C2 and C3 ar two different thicknesses of a solvent based acrylic adhesive containing Foral 85 resin. A mixture of 95.5 parts isooctyl acrylate and 4.5 parts of acrylic acid were polymerized using a free radical catalyst. A blend of 100 parts of polymerized adhesive in solvent is mixed with about 67 parts of Foral 85 resin. Example C2 is coated to a thickness of about 0.051 mm. Example C3 25 is coated to a thickness of about 0.127 mm. Comparative peel adhesion is listed in Table 2.

Table 2

90 Peel Adhesion (N/dm.)		
Example	Stainless Steel	Polycarbonate Plastic
2	79	88
3	118	113
C2	70	88
C3	125	113

#### Examples 4-15

These examples were made in substantially the same way as Example 1, but with different amounts of 45 Foral 85, photoinitiator and photocrosslinker at 2 mils and 5 mils thick. These are shown in Table 3. These samples were also tested accordingly and the results are listed in Table 4.

Table 3

Ex.	Foral 85	Irgacure 184	XL-330	Thickness
4	5	.25	.05	40 mm
5	5	.35	.05	40 mm
6	5	.25	.15	40 mm
7	5	.35	.15	40 mm
8	10	.30	.10	40 mm
9	10	.30	.10	40 mm
10	15	.25	.05	40 mm
11	15	.35	.05	40 mm
12	15	.25	.15	40 mm
13	15	.35	.15	40 mm
14	5	.25	.05	100 mm
15	15	.25	.05	100 mm

Table 4

	180° Peel Adhesion (24 Hr. Dwell at R.T., N/dm.)					
Ex.	Stainless Steel	Polypropylene	Polycarbonate	Shear at Room Temp. 1000 g (mins)		
4	67	41	55	130		
5	60	37	61	80		
6	41	30	53	3700		
7	42	34	51	2700		
8	63	42	60	110		
9	62	47	60	94		
10	60	78	51	6		
11	85	85	85	8		
12	73	47	68	68		
13	83	54	74	54		
14	60	55	72	70		
15	66	74	74	4		

## Examples 16-17

A premix was prepared using 95 parts isononyl acrylate, 5 parts acrylic acid, and 0.30 part IrgacureTM 184. This was partially polymerized by exposure to ultraviolet radiation to provide a coatable syrup. A blend of 100 parts of this partially polymerized premix, 10 parts Foral 85 resin, and 0.09 part inflaeureTM 184 was coated onto a polyethylene coated paper to a uniform thickness of about 0.05 mm as Example 16. Example 17 was prepared without tackfying resin. The surfaces of the polyethylene coated paper contacting the adhesive composition has low release coating the adhesive.

These composites were exposed to ultraviolet radiation to promote polymerization of the syrup. Comparative peel adhesion is listed in Table 5.

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Table 5

180 Peel Adhesion (N/dm.)		
Example	Polypropylene	ABS Plastic
16	94	94 48

#### Examples 18-19

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A premix was prepared using 90 parts butyl acrylate, 10 parts acrylic acid and 0.25 part Ingacure 164. 
This was partially polymertead by exposure to ultraviolet radiation to provide a coatable syrup. A blend of 
100 parts of this partially polymertead premix 10 parts Foral 85 reals in all 0.09 part Ingacure III 184 was 
cested onto a polyetylysiene coated paper to a uniform thickness of about 0.051 mm as Example 18. 
20 Example 19 was prepared without tackflying resin. The surfaces of the polyethylene coated paper 
contacting the adhesive composition has low release coatings on them.

contracting the subserve composites were exposed to ultraviolet radiation to promote polymerization of the syrup.

Comparative poel achesion is listed in Table 6.

Table 6

180* Peel Adhesion - 24 Hour Dwell (N/dm.)			
Example	Polypropylene	ABS Plastic	
18	57 20	83 70	

### Examples 20-23

These were made similar to Example 1 except with different amounts of Foral 85 and photocrosslinker. These differences are shown in Table 7 and test results are shown in Table 8.

Table 7

Example	Foral 85	XL-330
20	7	0.01
21	7	0.09
22	13	0.01
23	13	0.09

Table 8

180 Peel Adhesion (3 Day Dwell, Room Temp., N/dm.)				
Example	Polypropylene	ABS	Polyethylene	AI
20 21 22 23	103 87 85 100	161 145 178 176	91 76 83 87	218 198 167 169

# Examples 24-27 and Comparative Examples C24-C27

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These were made similar to Example 1 and Comparative Example 1 except with differing ratios of isocityl carylate and carylic acid. Example 27 was made using Foral 105, a pentaerithritol ester. These differences are shown in Table 3 and test results are shown in Table 10.

Table 9

Example	Acrytic Acid	Isooctyl Acrylate	Foral 85	Forel 105
24	1	99	10	_
C24	1	99	-	-
25	2	98	10	_
C25	2	98	-	
26	3	97	10	_
C26	3	97	-	_
27	6.5	93.5	_	10
C27	6.5	93.5	- 1	-

Table 10

180° Pee	180* Peel Adhesion (24 Hour Dwell, Room Temp., N/dm.)			
Example	Example Polypropylene			
24	47	61"		
C24	9	30		
25	42	56°		
C25	12	34		
	26 45			
C26	13	30		
27	43	77		
C27	19	53		

\* adhesive splits during test, indicating that adhesion to the substrate was so strong that the internal cohesion of the tape falled prior to fallure of the adhesive bond.

As can be clearly seen from the above data, adhesives of the invention provide improved peel adhesion over otherwise identical low acrylic acid formulations containing no tackfiller resin.

#### Claims

- 1. A tackfiled acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive comprising
- a) from about 85 parts to about 98 parts of an acrylic ultraviolet-radiation photopolymerized polymer
- of monomers containing: (f) from about 93 to about 99 parts of an alkyl acrylate monomer, the alkyl groups of which have an average of 4 to 12 carbon atoms, and
  - (ii) correspondingly, from about 7 parts to about 1 part of a monoethylenically unsaturated polar copolymerizable monomer,
- b) correspondingly, from about 15 parts to about 2 parts of a hydrogenated rosin ester tackifying
  - agent, and c) from about 0.01 part to about 1 part of a photoinitiator, based on the weight of the ultraviolet radiation polymerized polymer and the tackfrying resin.
- A tackfiled acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive according to claim 1
- wherein said tackifying agent comprises from about 5 to about 12 parts. 3. A tackified acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive according to claim t wherein said tacklifying agent is selected from the groups consisting of hydrogenated glycerine rosin esters,
- hydrogenated pentaerthritol esters, and glycol esters. 4. A tackfilled acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive composition according to claim 1 wherein said acrylic ultraviolet-radiation polymerized polymer comprises
  - a) from about 93 parts to about 99 parts of an alkyl acrylate ester selected from the group consisting of isooctyl acrylate, isononyl acrylate, decyl acrylate, dodecyl acrylate, butyl acrylate, ethyl-hexyl acrylate and hexyl acrylate,
  - b) correspondingly from about 7 parts to about 1 part of a monoethylenically unsaturated strongly polar copolymentzable monomer selected from the group consisting of acrylic acid, Itaconic acid, hydroxysikyl acrylates, cyanoalkyl acrylates, acrylamides or substituted acrylamides.
- 5. A tackfilled acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive according to claim 4 30 wherein said acrylic ultraviolet-radiation polymerized polymer comprises
  - a) from about 93 parts to about 99 parts isooctyl acrylate, and
  - b) correspondingly, from about 7 parts to about 1 part acrylic acid.
  - 6. A tackified acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive according to claim 1 wherein said acrylic ultraviolet-radiation polymerized polymer comprises
    - a) from about 95 to about 99 parts isooctyl acrylate, and
    - b) correspondingly from about 5 parts to about 1 part acrylic acid.
- 7. A tackified acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive according to claim 1 to further comprising from about 0.1 to about 1 part crosslinking agent.
  - 8. A tackified acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive according to claim 7
  - wherein said crosslinking agent is selected from the group consisting of triazines and multiacrylates. 9. A tackified acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive according to claim 1
- wherein said adhesive further comprises microspheres. 10. A pressure-sensitive adhesive tape comprising said tackfiled acrylic ultraviolet-radiation polymerized pressure-sensitive adhesive of claim 1 having at least one surface adhered to a flexible carrier web.

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		Relevant	CLASSIFICATION OF THE
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CATEGORY OF CITED DOCUMES: particularly relevant if taken alone particularly relevant if combined with and document of the same category sechological background com-written disclosure	F : mail or principle	ministrying the inve	